Filing Date: January 15, 2008

Title: Inhibiting Cav3 Isoforms and the 25B Splice Variants for the Diagnosis and Treatment of Cancer (Amended)

## REMARKS

This responds to the Final Office Action dated July 27, 2011.

Claims 7 and 10 are amended, claims 1-6 and 13-15 are withdrawn, claims 11-12 are cancelled, no claims are added; as a result, claims 1-10 and 13-15 are pending in this application.

Applicants thank the Examiner for noting the typographical errors in claims 7 and 10. The amendments to claims 7 and 10 render the objection of the claims moot. Thus, Applicants respectfully request withdrawal of the objection.

## The Rejection of Claims Under § 112

Claims 7-9 are rejected under 35 U.S.C. § 112, first paragraph, as allegedly lacking an adequate written description in the specification. Specifically, the Office Action contends that the instant specification does not contain a written description of the invention in such full, clear, concise, and exact terms or in sufficient detail that one skilled in the art can reasonably conclude that applicant had possession of the claimed invention at the time of filing.

Applicants respectfully disagree with and traverse this rejection. Applicants respectfully submit that the specification contains an adequate written description as follows.

Claim 7 currently recites "a method for inducing cytostasis comprising administering to a patient in need thereof a therapeutically effective amount T type calcium channel selective inhibitor so as to induce cytostasis in said patient." Support for claim 7 can be found in paragraph [0005] of the published application (Pub. No. 2008/0160009), which states the library of compounds that the present invention has developed can act cytostatically. In addition, paragraphs [0072] and [0073] and FIGS. 1A-1F of the present application teach that mibefradil can inhibit calcium entry in cancer cell lines. It is well known by one of skill in the art that inhibition of calcium entry can reversibly inhibit cell growth. As one of skill in the art would readily appreciate, reversible inhibition of cell growth is consistent with cytostasis rather than cytotoxicity. Such an inference would be made because preferred cytotoxic agents serve not in a

<sup>&</sup>lt;sup>1</sup> Whitfield JF, Boynton AL, MacManus JP, Sikorska M, Tsang BK. (1979) The regulation of cell proliferation by calcium and cyclic AMP. Mol Cell Biochem, 27, 152-179; Veigl ML, Sedwick WD, Vanaman TC. (1982) Calmodulin and Ca<sup>2+</sup> in normal and transformed cells. Federation Proceedings, 82, 2283-2288.

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reversible manner, but instead act to induce cell death.2 Therefore, one of skill in the art in possession of the present specification would understand that the figures and examples provided therein were aimed not at inducing cell death, but were aimed at reversible inhibition of cell growth, namely cytostasis, as presently recited in claim 7.

Claims 8 and 9 are dependent upon claim 7. Accordingly, claims 8 and 9 incorporate the limitations of claim 7. As such, Applicants respectfully submit that the features of claims 7-9 are fully supported by the application as originally filed. Applicants respectfully request withdrawal of this rejection.

Claims 7-9 are rejected under 35 U.S.C. § 112, first paragraph, as allegedly failing to comply with the enablement requirement. Specifically, the Office Action contends that the instant specification does not contain subject matter which is described in the specification in such a way as to enable one skilled in the art to which it pertains, or with which it is most nearly connected, to make and/or use the invention.

The Office Action states "Tolne of skill in the art cannot extrapolate the teachings of the specification to enable the claims because the claims are drawn to a method for inducing cytostasis comprising administering to a patient in need thereof a therapeutically effective amount of mibefradil so as to induce cytostasis in said patient; however, the specification has only presented data showing that mibefradil inhibits proliferation of prostate cancer cell lines in vitro; the specification presents no examples to demonstrate that mibefradil can induce cytostasis in cells or in a patient."

The Applicants respectfully disagree with and traverse this rejection. The Applicants respectfully submit that the specification adequately complies with the enablement requirement as follows.

As discussed above, claim 7 currently recites, among other things, "a method for inducing cytostasis...so as to induce cytostasis in said patient." For adequate enabling support the Applicants point to the teachings of paragraph [0005], which states the library of compounds that the present invention has developed can act cytostatically and, paragraphs [0072] and [0073] and FIGS. 1A-1F, which teach that mibefradil blocked the calcium entry from the extracellular medium that is necessary for cancer cell division and proliferation.

<sup>&</sup>lt;sup>2</sup>Blagosklonny MV (2005) Carcinogenesis, cancer therapy and chemoprevention. Cell Death Differ, 12, 592-602.

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First, the MPEP explicitly states that the question of compliance with the enablement requirement of 35 U.S.C. 112 first paragraph, "does not turn on whether an example is disclosed. An example may be 'working' or 'prophetic.'" The MPEP further explains that a working example can be based upon work performed, whereas a prophetic example describes an embodiment based on predicted results rather that the results actually obtained. All that is required is that there must be a correlation between the example contained in the specification and the claim. In fact, the Federal Circuit has reversed PTO decisions based on the erroneous finding that in vitro data did not support in vivo applications.4

Second, the MPEP explicitly states that because "the initial burden is on the examiner to give reasons for the lack of enablement, the examiner must also give reasons for a conclusion of lack of correlation for an in vitro or in vivo animal model." The Office Action states that "the claims are drawn to a method for inducing cytostasis comprising administering to a patient in need thereof a therapeutically effective amount of mibefradil so as to induce cytostasis in said patient; however, the specification has only presented data showing that mibefradil inhibits proliferation of prostate cancer cell lines in vitro." As such, Applicants respectfully assert that the instant 35 U.S.C. § 112, first paragraph rejection is improper because the Office Action has failed to give reasons for a conclusion of lack of correlation between the in vitro data and an in vivo.

Finally, the MPEP states that a specification need not contain a working example if the invention is disclosed in a manner such that one skilled in the art will be able to practice it without an undue amount of experimentation. It is well settled law that at times, even when the amount of experimentation required to practice the scope of the claimed invention might have been extensive, the experimentation is still routine if the necessary techniques are well known to those skilled in the art. In the specific instance of mibefradil, it is well known in the art that

<sup>&</sup>lt;sup>3</sup> U.S. Pat. & Trademark Office, U.S. Dep't of Commerce, Manual of Patent Examining Procedure § 2164.02 (8th ed., 2nd rev. 2004) [hereinafter MPEP].

See In re Brana, 51 F.3d 1560, 1566, 34 USPO2d 1436, 1441 (Fed. Cir. 1995).

<sup>5</sup> MPEP 8 2164.02.

<sup>6</sup> Id.: In re Borkowski, 422 F.2d 904, 908, 164 USPO 642, 645 (CCPA 1970).

<sup>&</sup>lt;sup>7</sup> Ex parte Kubin, Appeal 2004-0819, 2007 WL 2070495 (B.P.A.I. May 31, 2007).

certain doses are recommended for administration to a patient in need thereof.8 As such, the Applicants respectfully assert that although one of skill in the art in possession of the present disclosure may need to perform routine experimentation to obtain the appropriate dose of mibefradil for therapeutic effect in a patient, such routine experimentation is not extensive, and does not rise to the level of impermissible undue experimentation.

Claims 8 and 9 are dependent upon claim 7. Accordingly, claims 8 and 9 incorporate the limitations of claim 7. As such Applicants respectfully submit that the features of claims 7-9 are enabled and comply with 112(1). Thus, Applicants respectfully request withdrawal of this rejection.

## The Rejection of Claims Under § 103

Claim 10 remains rejected under 35 U.S.C. 103 (a) as allegedly being unpatentable over Bertolesi et al. (Mol. Pharmacol. 62:210-219, 2002, hereafter Bertolesi) in view of Gray et al. (U.S. Patent Number 6413967, hereafter '967).

Applicants respectfully traverse the rejections and request reconsideration and withdrawal of the rejection.

The U.S. Supreme Court's decision of KSR v. Teleflex<sup>9</sup> provided a multi-prong test to evaluate obviousness. This multi-prong test set forth a list of requirements to support a conclusion of obviousness, including-among other things-a finding that all the claimed elements were known in the prior art. 10 Applicants maintain that the limitation of inducing cytostasis in a patient, as claimed in currently amended claim 10, is not disclosed in the cited art.

Applicants cannot find in the cited portions of Bertolesi, Gray or the Office Action's reasoning a disclosure of inducing cytostasis in a patient. In fact, Bertolesi appears to expressly teach away from this. Bertolesi explicitly rejects the hypothesis that T type calcium blockers inhibit proliferation by cell cycle arrest and cytostasis (page 214 section beginning in left column titled "Cytostatic or Cytotoxic Effects of Pimoxzide and Mibefradil") (emphasis added).

<sup>&</sup>lt;sup>8</sup> Welker HA (1998) Single- and multiple-dose mibefradil pharmacokinetics in normal and hypertensive subjects. J Pharm Pharmacol, 50, 983-7,

<sup>&</sup>lt;sup>9</sup> KSR International Co. v. Teleflex Inc., 127 S. Ct, 1727, 82 U.S.P.Q.2d 1385 (2007).

<sup>&</sup>lt;sup>10</sup> See also, Manual of Patent Examining Procedure §§ 706.02(j), 2143(A) (2008); MPEP § 2142 (2006) (citing In re Vaeck, 947 F.2d, 488 (Fed. Cir. 1991)). Cited approvingly in Exparte Wen Wen and Patricia NG at 7; Appeal No. 2009-000776; decided September 25, 2009.

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Bertolesi interprets the results to indicate that these drugs block proliferation by inducing cell death. Conversely, the major premise of the presently claimed invention is that cytostasis is induced by inhibition by T type calcium blockers and thereby inducing a cell cycle blockade. As previously pointed out by the Applicants, cytostatic and cytotoxic mechanisms are fundamentally inapposite. For example, cytotoxic drugs cause collateral damage to normal, healthy tissues, which bring dose and schedule limiting toxicities. On the other hand, cytostatic drugs can be administered to patients chronically. As implicitly acknowledged by Bertolesi by investigating cytostatic or cytotoxic effects rather than cytostatic and cytotoxic effects, cytotoxicity and cytostasis is mutually exclusive. In sum, Applicants respectfully submit that Bertolesi not only fails to disclose or suggest inducing cytostasis in a patient, Bertolesi actually teaches away from the subject matter of claim 10, as currently amended.

The Applicant respectfully asserts that Gray also fails to disclose or suggest inducing cytostasis or inducing cytostasis in a patient. As such, the Applicants assert that Gray does not remedy the deficiencies of Bertolesi.

As the 35 USC § 103 rejection of the Office Action has been obviated by the amendments to independent claim 10, claim 10 is presently believed to be in allowable condition. Reconsideration and withdrawal of the rejection of claim 10 is respectfully requested.

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## CONCLUSION

Applicant respectfully submits that the claims are in condition for allowance, and notification to that effect is earnestly requested. The Examiner is invited to telephone Applicant's representative at (612) 373-6905 to facilitate prosecution of this application.

If necessary, please charge any additional fees or deficiencies, or credit any overpayments to Deposit Account No. 19-0743.

Respectfully submitted,

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By / Monique U. Pardyk Shonka

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